```
import numpy as np
          import matplotlib.pyplot as plt
          from tensorflow.keras.models import load model
          from sklearn.model selection import StratifiedKFold, train test split
          from sklearn.metrics import accuracy score, fl score, precision score, recall score, roc auc score, average precision score
          from tensorflow.keras.utils import to categorical
          import tensorflow as tf
          import pandas as pd
          import model get
          import pickle as pkl
In [2]:
          import seaborn as sns
          from sklearn.metrics import roc curve, auc, precision recall curve
          physical devices = tf. config. experimental. list physical devices ('GPU')
          if len(physical devices) > 0:
               tf. config. experimental. set memory growth (physical devices [0], False)
          def loadGlove(inputpath, outputpath=""):
               data list = []
               wordEmb = \{\}
              with open (input path) as f:
                   for line in f:
                       11 = line. strip(). split(',')
                       11[0] = str(int(float(11[0])))
                       data list. append (11)
                       11 \text{ new} = \lceil \text{float}(i) \text{ for } i \text{ in } 11 \rceil
                       emb = np. array (11 new[1:], dtype="float32")
                       wordEmb[str(int(11 new[0]))] = emb
              if outputpath != "":
                   with open (outputpath) as f:
                       for data in data list:
                           f. writelines(' '. join(data))
              return wordEmb
          def plotPrecisionRecallCurve(estimators, labels, xtests, ytests, flnm, icol=1):
               colors = sns. color palette ("hus1", len (estimators)) # Use a color palette from seaborn
```

```
indx = 0
    plt. figure (figsize=(10, 6)) # Increase figure size
   plt. plot([0, 1], [0, 1], 'k--')
   for estimator, color in zip(estimators, colors):
        if len(ytests[indx]. shape) == 2:
           pre, rec, = precision recall curve(ytests[indx][:, icol], estimator.predict(xtests[indx])[:, icol], pos label=icol)
       else:
            pre, rec, = precision recall curve(ytests[indx], estimator.predict proba(xtests[indx])[:, icol], pos label=icol)
       plt. plot(rec. pre. label=labels[indx] + ' (PR AUC: %s \u0000B1 0.001)' % (np. round(auc(rec. pre), 3)), color=color)
       indx += 1
   plt. xlabel ('Recall', fontsize=14) # Increase font size
    plt. vlabel ('Precision', fontsize=14) # Increase font size
    plt. legend (loc='best', fontsize=12) # Increase font size
    plt. savefig (flnm)
def plotRocCurve(estimators, labels, xtests, ytests, flnm, icol=1):
    colors = sns.color palette ("husl", len (estimators)) # Use a color palette from seaborn
    indx = 0
   plt. figure (figsize=(10, 6)) # Increase figure size
   plt. plot([0, 1], [0, 1], 'k--')
   for estimator, color in zip(estimators, colors):
        if len(ytests[indx]. shape) == 2:
           fprs, tprs, = roc curve(ytests[indx][:, icol], estimator.predict(xtests[indx])[:, icol])
            fprs, tprs, = roc curve(ytests[indx], estimator.predict proba(xtests[indx])[:, icol])
       plt.plot(fprs, tprs, label=labels[indx] + '(ROC AUC: %s \u00B1 0.001)' % (np.round(auc(fprs, tprs), 3)), color=color)
       indx += 1
    plt. xlabel ('False positive rate', fontsize=14) # Increase font size
   plt.ylabel('True positive rate', fontsize=14) # Increase font size
    plt. legend (loc='best', fontsize=12) # Increase font size
    plt. savefig (flnm)
```

```
import random
import os
seed = 123
random. seed(seed)
os. environ['PYTHONHASHSEED']=str(seed)
np. random. seed(seed)
tf. random. set_seed(seed)
```

```
def f1 metric(y true, y pred):
    true positives = tf. keras. backend. sum(tf. keras. backend. round(tf. keras. backend. clip(y true * y pred, 0, 1)))
    possible positives = tf. keras. backend. sum(tf. keras. backend. round(tf. keras. backend. clip(y true, 0, 1)))
    predicted positives = tf. keras. backend. sum(tf. keras. backend. round(tf. keras. backend. clip(y pred, 0, 1)))
    precision = true positives / (predicted positives + tf. keras. backend. epsilon())
    recall = true positives / (possible positives + tf. keras. backend. epsilon())
    f1 val = 2 * (precision * recall) / (precision + recall + tf. keras. backend. epsilon())
    return fl val
early stopping = tf. keras. callbacks. EarlyStopping(
    monitor='val fl metric', min delta=0.0001,
    patience=10, verbose=0, mode='max')
callbacks = [early stopping]
dataset = 'ELEVATION'
num classes = 2
flpath = 'data/'
retrain = False
```

cross-validation

Crispr_IP

```
# Initialize the cross-validation splitter
cv = StratifiedKFold(n_splits=5, random_state=42, shuffle=True)

print('crispr_ip_model')
encoder_shape = (23, 6)
seg_len, coding_dim = encoder_shape

open_name = 'encoded6x23' + dataset + '.pkl'

print('load_data!')
print(open_name)

loaddata = pkl.load(
open(flpath + open_name, 'rb'),
```

```
encoding='latin1'
# Prepare data for cross-validation
X = np. array(loaddata. images)
v = loaddata, target
eval fun names = ['Accuracy', 'F1 score', 'Precision', 'Recall', 'ROC AUC', 'PR AUC']
eval funs = [accuracy score, fl score, precision score, recall score, roc auc score, average precision score]
eval fun types = [True, True, True, True, False, False]
# Initialize the result dict
results1 = {name: [] for name in eval fun names}
# Load the pre-trained model
model path = 'saved model/'+dataset+'crispr ip. h5'
crispr ip model = load model(model path, custom objects={'fl metric': fl metric})
# Perform cross-validation
for train index, test index in cv. split(X, y):
   X train full, X test = X[train index], X[test index]
    v train full, v test = v[train index], v[test index]
    # Create a validation set from the full training data
   X train, X val, y train, y val = train test split(
        X train full,
        y train full,
        stratify=pd. Series(y train full),
        test size=0.2,
        shuffle=True,
        random state=42)
    # Transform the data
    X train, X testl, y train, y testl, X val, y val, input shape = model get.transformIO(
        X train, X test, y train, y test, X val, y val, seg len, coding dim, num classes
    # Calculate metrics
   y score = crispr ip model.predict(X test1)
    v pred = np. argmax(v score, axis=1)
    y score = y score[:, 1]
    y \text{ test} = \text{np. argmax}(y \text{ test1}, \text{ axis}=1)
    # Store the metrics
    for index f, function in enumerate (eval funs):
```

```
if eval fun types[index f]:
              score = np. round(function(y test, y pred), 4)
         else:
              score = np. round(function(y test, y score), 4)
         results1[eval fun names[index f]].append(score)
 # Calculate the average of metrics
 avg results1 = {name: np. mean(scores) for name, scores in results1. items()}
 # Print the average metrics
 for name, avg score in avg results1. items():
     print (\{:\langle 15\rangle\} \{:\rangle 15\}). format (name, avg score))
crispr ip model
 load data!
encoded6x23ELEVATION.pk1
xtrain shape: (136916, 1, 23, 6)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 6)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 6)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136917, 1, 23, 6)
136917 train samples
42786 test samples
34230 val samples
xtrain shape: (136917, 1, 23, 6)
136917 train samples
42786 test samples
34230 val samples
Accuracy
                        0.04122
                        0.00046
F1 score
                         0.0002
Precision
Recal1
                0.94000000000000001
ROC AUC
                0.5332799999999999
PR AUC
                         0.0036
cnn std
```

```
# Initialize the cross-validation splitter
cv = StratifiedKFold(n splits=5, random state=42, shuffle=True)
print('cnn std')
encoder shape = (23, 4)
seg len, coding dim = encoder shape
open name = 'encoded4x23' + dataset + '.pk1'
print('load data!')
print(open name)
loaddata = pkl. load(
    open(flpath + open name, 'rb'),
    encoding='latin1'
# Prepare data for cross-validation
X = np. array (loaddata. images)
y = loaddata. target
eval fun names = ['Accuracy', 'F1 score', 'Precision', 'Recall', 'ROC AUC', 'PR AUC']
eval funs = [accuracy score, fl score, precision score, recall score, roc auc score, average precision score]
eval fun types = [True, True, True, True, False, False]
# Initialize the result dict
results2 = {name: [] for name in eval fun names}
# Load the pre-trained model
model path = 'saved model/'+dataset+'cnn std.h5'
cnn std model = load model(model path, custom objects={'fl metric': fl metric})
# Perform cross-validation
for train index, test index in cv. split(X, y):
   X train full, X test = X[train index], X[test index]
    y train full, y test = y[train index], y[test index]
    # Create a validation set from the full training data
    X train, X val, y train, y val = train test split(
        X train full,
        y_train_full,
```

```
cross validation
         stratify=pd. Series(y train full),
         test size=0.2,
         shuffle=True,
         random state=42)
    # Transform the data
    X train, X test2, y train, y test2, X val, y val, input shape = model get.cnn std transformIO(
         X train, X test, y train, y test, X val, y val, seg len, coding dim, num classes
     # Calculate metrics
    y score = cnn std model.predict(X test2)
    y pred = np. argmax(y score, axis=1)
    y score = y score[:, 1]
    y test = np. argmax(y test2, axis=1)
    # Store the metrics
    for index f, function in enumerate (eval funs):
         if eval fun types [index f]:
             score = np. round(function(y test, y pred), 4)
         else:
             score = np. round(function(y test, y score), 4)
         results2[eval fun names[index f]]. append(score)
# Calculate the average of metrics
avg results2 = {name: np. mean(scores) for name, scores in results2. items()}
# Print the average metrics
for name, avg score in avg results2.items():
    print (\{:\langle 15\}\} \{:\rangle 15\}). format (name, avg score)
cnn std
```

```
load data!
encoded4x23ELEVATION.pk1
xtrain shape: (136916, 1, 23, 4)
136916 train samples
42787 test samples
34230 xval samples
xtrain shape: (136916, 1, 23, 4)
136916 train samples
42787 test samples
34230 xval samples
xtrain shape: (136916, 1, 23, 4)
136916 train samples
42787 test samples
34230 xval samples
```

```
xtrain shape: (136917, 1, 23, 4)
136917 train samples
42786 test samples
34230 xval samples
xtrain shape: (136917, 1, 23, 4)
136917 train samples
42786 test samples
34230 xval samples
                        0.0002
Accuracy
F1 score
                        0.0005
Precision
                        0.0002
                           1.0
Recal1
ROC AUC
                           0.5
PR AUC
                        0.0002
```

crisprDNT

```
from model get import PositionalEncoding
from keras multi head import MultiHeadAttention
# Initialize the cross-validation splitter
cv = StratifiedKFold(n splits=5, random state=42, shuffle=True)
print('crisprDNT')
encoder shape = (23, 14)
seg len, coding dim = encoder shape
open name = 'encodedmismatchtype14x23' + dataset + '.pk1'
print('load data!')
print(open name)
loaddata = pkl.load(
    open(flpath + open name, 'rb'),
    encoding='latin1'
# Prepare data for cross-validation
X = np. array(loaddata. images)
y = loaddata. target
eval fun names = ['Accuracy', 'F1 score', 'Precision', 'Recall', 'ROC AUC', 'PR AUC']
eval funs = [accuracy score, fl score, precision score, recall score, roc auc score, average precision score]
```

```
eval fun types = [True, True, True, True, False, False]
# Initialize the result dict
results3 = {name: [] for name in eval fun names}
# Load the pre-trained model
model path = 'saved model/'+dataset+'crisprDNT.h5'
crisprDNT = load model (model path, custom objects={'PositionalEncoding': PositionalEncoding,'MultiHeadAttention': MultiHeadAttention,
# Perform cross-validation
for train index, test index in cv. split(X, y):
    X train full, X test = X[train index], X[test index]
    v train full, v test = v[train index], v[test index]
    # Create a validation set from the full training data
    X train, X val, y train, y val = train test split(
        X train full,
        y train full,
        stratify=pd. Series(y train full),
        test size=0.2,
        shuffle=True,
        random state=42)
    # Transform the data
    X train, X test3, y train, y test3, X val, y val, input shape = model get.transformIO(
        X train, X test, y train, y test, X val, y val, seg len, coding dim, num classes
    # Calculate metrics
    y score = crisprDNT. predict(X test3)
    y pred = np. argmax(y score, axis=1)
    y score = y score[:, 1]
    y test = np. argmax(y test3, axis=1)
    # Store the metrics
    for index f, function in enumerate (eval funs):
        if eval fun types [index f]:
            score = np. round(function(y test, y pred), 4)
            score = np. round(function(y test, y score), 4)
        results3[eval fun names[index f]].append(score)
# Calculate the average of metrics
avg results3 = {name: np. mean(scores) for name, scores in results3. items()}
```

```
# Print the average metrics
 for name, avg score in avg results3. items():
     print (\{:\langle 15\rangle\} \{:\rangle 15\}). format (name, avg score))
crisprDNT
load data!
encodedmismatchtype14x23ELEVATION.pk1
WARNING: tensorflow: Layer 1stm 1 will not use cuDNN kernel since it doesn't meet the cuDNN kernel criteria. It will use generic GPU ke
rnel as fallback when running on GPU
WARNING: tensorflow: Layer 1stm 1 will not use cuDNN kernel since it doesn't meet the cuDNN kernel criteria. It will use generic GPU ke
rnel as fallback when running on GPU
WARNING: tensorflow: Layer 1stm 1 will not use cuDNN kernel since it doesn't meet the cuDNN kernel criteria. It will use generic GPU ke
rnel as fallback when running on GPU
xtrain shape: (136916, 1, 23, 14)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 14)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 14)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136917, 1, 23, 14)
136917 train samples
42786 test samples
34230 val samples
xtrain shape: (136917, 1, 23, 14)
136917 train samples
42786 test samples
34230 val samples
                0.9996400000000001
Accuracy
F1 score
                0.4918600000000001
Precision
                         0.3722
Recall
                           0.74
ROC AUC
                        0.99512
PR AUC
                        0.68282
crisprNet
 # Initialize the cross-validation splitter
 cv = StratifiedKFold(n splits=5, random state=42, shuffle=True)
 print('CRISPR Net')
```

```
encoder shape = (23, 7)
seg len, coding dim = encoder shape
open name = 'encoded7x23' + dataset + '.pk1'
print('load data!')
print(open name)
loaddata = pkl. load(
    open(flpath + open name, 'rb'),
    encoding='latin1'
# Prepare data for cross-validation
X = np. array (loaddata. images)
y = loaddata. target
eval fun names = ['Accuracy', 'F1 score', 'Precision', 'Recall', 'ROC AUC', 'PR AUC']
eval funs = [accuracy score, fl score, precision score, recall score, roc auc score, average precision score]
eval fun types = [True, True, True, True, False, False]
# Initialize the result dict
results4 = {name: [] for name in eval fun names}
# Load the pre-trained model
model path = 'saved model/'+dataset+'CRISPR Net.h5'
CRISPR Net model = load model (model path, custom objects={'f1 metric': f1 metric})
# Perform cross-validation
for train index, test index in cv. split(X, y):
   X train full, X test = X[train index], X[test index]
    y train full, y test = y[train index], y[test index]
    # Create a validation set from the full training data
   X train, X val, y train, y val = train test split(
        X train full,
        y_train_full,
        stratify=pd. Series(y train full),
        test size=0.2,
        shuffle=True,
        random state=42)
    # Transform the data
```

```
cross validation
    X_train, X_test4, y_train, y_test4, X val, y val, input shape = model get. CRISPR Net transformIO(
         X train, X test, y train, y test, X val, y val, seg len, coding dim, num classes
     # Calculate metrics
    y score = CRISPR Net model.predict(X test4)
    y pred = np. argmax(y score, axis=1)
     y score = y score[:, 1]
    y test = np. argmax(y test4, axis=1)
     # Store the metrics
    for index f, function in enumerate (eval funs):
         if eval fun types [index f]:
             score = np. round(function(y test, y pred), 4)
         else:
             score = np. round (function (y test, y score), 4)
         results4 [eval fun names [index f]]. append (score)
# Calculate the average of metrics
avg results4 = {name: np. mean(scores) for name, scores in results4. items()}
# Print the average metrics
for name, avg score in avg results4. items():
    print (\{:\langle 15\rangle\} \{:\rangle 15\}). format (name, avg score))
CRISPR Net
load data!
encoded7x23ELEVATION.pk1
xtrain shape: (136916, 1, 23, 7)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 7)
136916 train samples
42787 test samples
34230 val samples
xtrain shape: (136916, 1, 23, 7)
136916 train samples
42787 test samples
```

136917 train samples

34230 val samples

136917 train samples 42786 test samples 34230 val samples

xtrain shape: (136917, 1, 23, 7)

xtrain shape: (136917, 1, 23, 7)

```
34230 val samples
               0.9998000000000001
Accuracy
F1 score
                       0.63226
Precision
                       0.53888
                           0.78
Recall
ROC AUC
                        0.99116
PR AUC
                       0.71818
cnnCRISPR
```

```
# Initialize the cross-validation splitter
cv = StratifiedKFold(n splits=5, random state=42, shuffle=True)
print('cnn crispr model')
print("GloVe model loaded")
VOCAB SIZE = 16 # 4**3
EMBED SIZE = 100
glove inputpath = "data/keras GloVeVec" + dataset + " 5 100 10000.csv"
# load GloVe model
model glove = loadGlove(glove inputpath)
embedding weights = np. zeros((VOCAB SIZE, EMBED SIZE))
for i in range (VOCAB SIZE):
    embedding weights[i, :] = model glove[str(i)]
open name = 'encoded CnnCrispr' + dataset + '.pkl'
print('load data!')
print(open name)
loaddata = pkl. load(
    open(flpath + open name, 'rb'),
    encoding='latin1'
# Prepare data for cross-validation
X = np. array (loaddata. images)
y = loaddata. target
eval fun names = ['Accuracy', 'F1 score', 'Precision', 'Recall', 'ROC AUC', 'PR AUC']
eval funs = [accuracy score, f1 score, precision score, recall score, roc auc score, average precision score]
eval fun types = [True, True, True, True, False, False]
# Initialize the result dict
results5 = {name: [] for name in eval fun names}
```

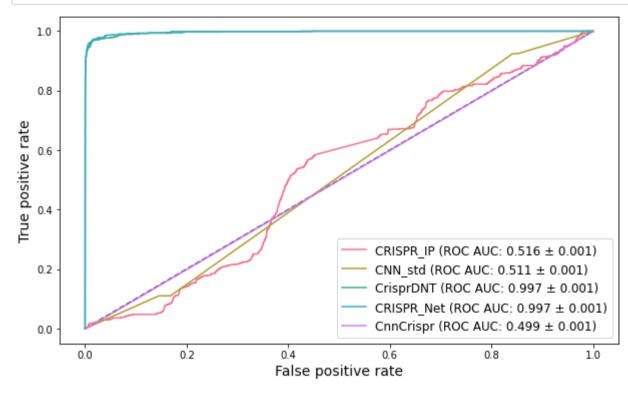
```
# Load the pre-trained model
model path = 'saved model/'+dataset+'CnnCrispr.h5'
CnnCrispr model = load model(model path, custom objects={'f1 metric': f1 metric})
# Perform cross-validation
for train index, test index in cv. split(X, y):
    X train full, X test = X[train index], X[test index]
    v train full, v test = v[train index], v[test index]
    # Create a validation set from the full training data
    X train, X val, y train, y val = train test split(
        X train full,
        y train full,
        stratify=pd. Series(y train full),
        test size=0.2,
        shuffle=True,
        random state=42)
    # Transform the data
    X train, X test5, y train, y test5, X val, y val = model get.offt transformIO(
        X train, X test, y train, y test, X val, y val, num classes
    # Calculate metrics
    y score = CnnCrispr model.predict(X test5)
    y pred = np. argmax(y score, axis=1)
    y score = y score[:, 1]
    y \text{ test} = \text{np. argmax}(y \text{ test5}, \text{ axis}=1)
    # Store the metrics
    for index f, function in enumerate (eval funs):
        if eval fun types[index f]:
            score = np. round(function(y test, y pred), 4)
        else:
            score = np. round(function(y test, y score), 4)
        results5[eval fun names[index f]]. append(score)
# Calculate the average of metrics
avg results5 = {name: np. mean(scores) for name, scores in results5. items()}
# Print the average metrics
for name, avg score in avg results5. items():
    print (\{:\langle 15\} \{:\rangle 15\}). format (name, avg score))
```

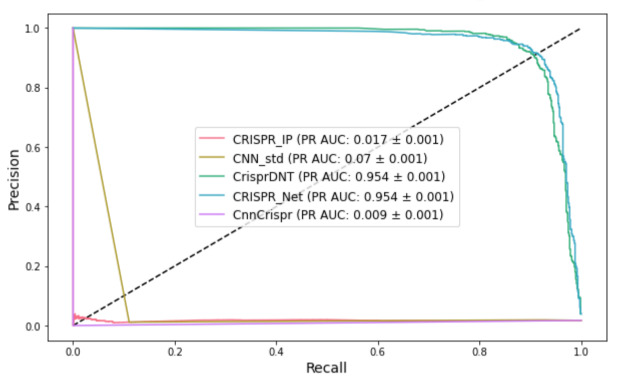
```
cnn crispr model
GloVe model loaded
FileNotFoundError
                                           Traceback (most recent call last)
<ipython-input-95-9ba069ead4ee> in <module>
     8 glove inputpath = "data/keras GloVeVec" + dataset + " 5 100 10000.csv"
      9 # load GloVe model
---> 10 model glove = loadGlove (glove inputpath)
     11 embedding weights = np. zeros ((VOCAB SIZE, EMBED SIZE))
     12 for i in range (VOCAB SIZE):
<ipython-input-2-2e43998b75d0> in loadGlove(inputpath, outputpath)
           data list = []
           wordEmb = \{\}
     10
---> 11
             with open (input path) as f:
     12
               for line in f:
     13
                   11 = line. strip(). split(',')
FileNotFoundError: [Errno 2] No such file or directory: 'data/keras GloVeVec ELEVATION 5 100 10000.csv'
```

Plot

```
In [88]: # models = [ crispr_ip_model, cnn_std_model, crisprDNT, CRISPR_Net_model, CnnCrispr_model]
# labels = ['CRISPR_IP', 'CNN_std', 'CrisprDNT', 'CRISPR_Net', 'CnnCrispr']
# xtests = [X_testl, X_test2, X_test3, X_test4, X_test5]
# ytests = [y_testl, y_test2, y_test3, y_test4, y_test5]
# roc_name = 'fig/ROC_5fold_' + dataset + '.png'
# pr_name = 'fig/PR_5fold_' + dataset + '.png'
# plotRocCurve(models, labels, xtests, ytests, roc_name)
# plotPrecisionRecallCurve(models, labels, xtests, ytests, pr_name)
models = [ crispr_ip_model, cnn_std_model, crisprDNT, CRISPR_Net_model]
labels = ['CRISPR_IP', 'CNN_std', 'CrisprDNT', 'CRISPR_Net']
xtests = [X_testl, X_test2, X_test3, X_test4]
ytests = [y_test1, y_test2, y_test3, y_test4]
```

```
roc_name = 'fig/ROC_5fold_' + dataset + '.png'
pr_name = 'fig/PR_5fold_' + dataset + '.png'
plotRocCurve(models, labels, xtests, ytests, roc_name)
plotPrecisionRecallCurve(models, labels, xtests, ytests, pr_name)
```





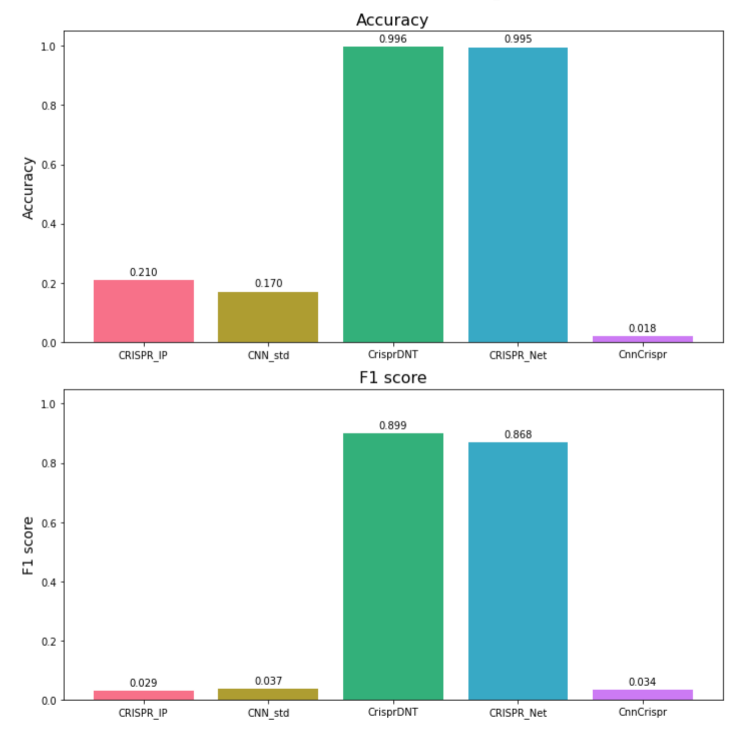
```
In [89]:
           # # Extract the metrics for each model
           # accuracy scores1 = avg results1['Accuracy']
           # accuracy scores2 = avg results2['Accuracy'
           # accuracy scores3 = avg results3['Accuracy'
           # accuracy scores4 = avg results4 ['Accuracy'
           # accuracy scores5 = avg results5['Accuracy'
           # fl scores1 = avg results1['Fl score']
           # fl scores2 = avg results2['Fl score']
           # f1 scores3 = avg results3['F1 score']
           # fl scores4 = avg results4['Fl score']
           # f1 scores5 = avg results5['F1 score']
           # precision scores1 = avg results1['Precision']
           # precision scores2 = avg results2['Precision'
           # precision scores3 = avg results3['Precision'
           # precision scores4 = avg results4['Precision'
           # precision scores5 = avg results5['Precision']
           # recall scores1 = avg results1['Recall']
           # recall scores2 = avg results2['Recall']
           # recall scores3 = avg results3['Recall']
           # recall scores4 = avg results4['Recall']
           # recall scores5 = avg results5['Recall']
           # # Create dictionaries for each metric that contains the scores for all models
           # accuracy scores = [accuracy scores1, accuracy scores2, accuracy scores3, accuracy scores4, accuracy scores5]
           # f1 scores = [f1 scores1, f1 scores2, f1 scores3, f1 scores4, f1 scores5]
           # precision scores = [precision scores1, precision scores2, precision scores3, precision scores4, precision scores5]
           # recall scores = [recall scores1, recall scores2, recall scores3, recall scores4, recall scores5]
           # metrics = {
```

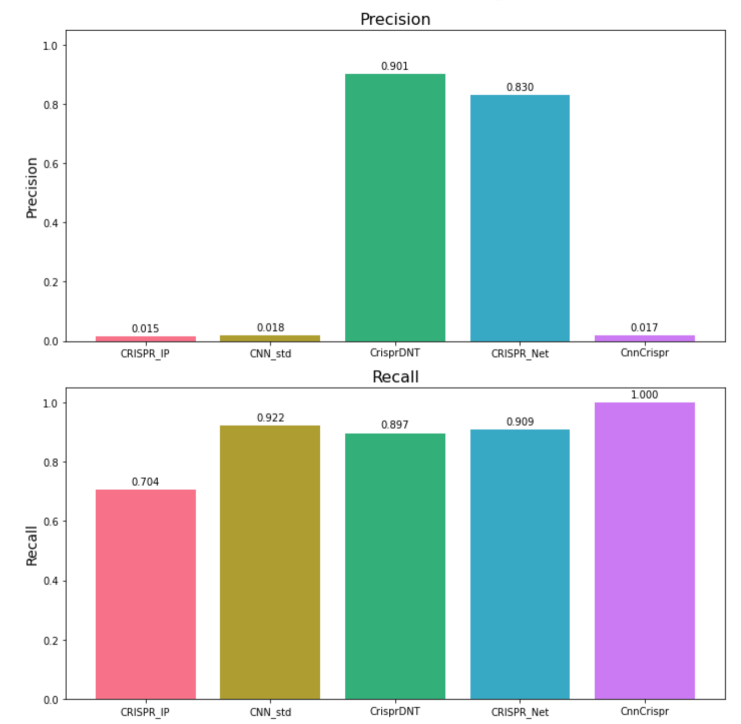
```
'Accuracy': accuracy scores,
     'F1 score': f1 scores,
     'Precision': precision scores,
      'Recall': recall scores
# }
# labels = ['CRISPR IP', 'CNN std', 'CrisprDNT', 'CRISPR Net', 'CnnCrispr']
# plot metrics(metrics, labels, 'fig/5-fold metrics plot SITE.png')
# Extract the metrics for each model
accuracy scores1 = avg results1['Accuracy']
accuracy scores2 = avg results2['Accuracy']
accuracy scores3 = avg results3['Accuracy']
accuracy scores4 = avg results4['Accuracy']
fl scores1 = avg results1['Fl score']
f1 scores2 = avg results2['F1 score']
f1 scores3 = avg results3['F1 score']
f1 scores4 = avg results4['F1 score']
precision scores1 = avg results1['Precision']
precision scores2 = avg results2['Precision'
precision scores3 = avg results3['Precision'
precision scores4 = avg results4['Precision']
recall scores1 = avg results1['Recall']
recall scores2 = avg results2['Recall']
recall scores3 = avg results3['Recall']
recall scores4 = avg results4['Recall']
# Create dictionaries for each metric that contains the scores for all models
accuracy scores = [accuracy scores1, accuracy scores2, accuracy scores3, accuracy scores4]
fl scores = [fl scores1, fl scores2, fl scores3, fl scores4]
precision scores = [precision scores1, precision scores2, precision scores3, precision scores4]
recall scores = [recall scores1, recall scores2, recall scores3, recall scores4]
metrics = {
    'Accuracy': accuracy scores,
    'F1 score': f1 scores,
    'Precision': precision scores,
```

```
'Recall': recall_scores
}

labels = ['CRISPR_IP', 'CNN_std', 'CrisprDNT', 'CRISPR_Net']

plot_metrics(metrics, labels, 'fig/5-fold_metrics_plot_ELEVATION.png')
```





In []: